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**Bacillus anthracis** lethal toxin induces TNF-α–independent hypoxia-mediated toxicity in mice

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**Bacillus anthracis** lethal toxin (LT) is the major virulence factor of anthrax and reproduces most of the laboratory manifestations of the disease in animals. We studied LT toxicity in BALB/cj and C57BL/6J mice. BALB/cj mice became terminally ill earlier and with higher frequency than C57BL/6J mice. Timed histopathological analysis identified bone marrow, spleen, and liver as major affected organs in both mouse strains. LT induced extensive hypoxia. Crisis was due to extensive liver necrosis accompanied by pleural edema. There was no evidence of disseminated intravascular coagulation or renal dysfunction. Instead, analyses revealed hepatic dysfunction, hypoalbuminemia, and vascular/oxygenation insufficiency. Of 50 cytokines analyzed, BALB/cj mice showed rapid but transitory increases in specific factors including KC, MCP-1/JE, IL-6, MIP-2, G-CSF, GM-CSF, eotaxin, FasL, and IL-1β. No changes in TNF-α occurred. The C57BL/6J mice did not mount a similar cytokine response. These factors were not induced in vitro by LT treatment of toxin-sensitive macrophages. The evidence presented shows that LT kills mice through a TNF-α–independent, FasL-independent, noninflammatory mechanism that involves hypoxic tissue injury but does not require macrophage sensitivity to toxin.


**Introduction**

Anthrax, the disease caused by *Bacillus anthracis*, is a worldwide bioterrorism concern. Anthrax toxin, the major virulence factor of this organism, consists of three polypeptides: edema factor (EF), lethal factor (LF), and protective antigen (PA). EF (an adenyl cyclase) and LF (a protease) are toxic only when combined with PA. PA is required for binding and translocation of these enzymatic moieties into target cells (for review, see ref. 1). Lethal toxin (LT), the combination of LF and PA, is sufficient to induce many of the laboratory manifestations of anthrax disease in animal models (2–6). Early studies suggested that LT killed animals by inducing nonspecific shocklike manifestations and vascular leakage (2, 7). Pulmonary edema has also been associated with many models of LT-mediated lethality, including the uniquely rapid death of Fischer 344 rats (8). Humans and primates infected by aerosol exposure to spores present pleural effusions as the most common symptom of disease (9–14).

Comparison of mutated *B. anthracis* strains lacking individual toxin components proved that LT is the major virulence factor in mice (15, 16). The discovery that inbred mouse strains differed in their relative susceptibility to LT, and that this correlated with the LT sensitivity of their macrophages, implicated this cell in the pathogenesis of LT (17). Later, macrophage depletion experiments in BALB/c mice led Hanna et al. to conclude that macrophage sensitivity was a requirement for LT-mediated toxicity in mice (18). Reports that very low doses of LT induce sensitive macrophages to produce TNF-α and IL-1β in vitro, along with the ability of an IL-1 antagonist or a mixture of antisera to TNF-α and IL-1β to protect mice from LT challenge, led to the suggestion of a macrophage-dependent endotoxic shock–like death in anthrax (18). This view of macrophage-dependent pathogenesis has been widely cited and has influenced many studies being initiated to identify therapies for treatment of anthrax infections. Thus, many investigators are focusing on anticytokine therapies similar to those used for sepsis. However, in recent years various laboratories have been unable to induce production of inflammatory cytokines by LT-treated toxin-sensitive macrophages in vitro at the RNA or protein level (19). In fact, the ability of macrophages to produce these factors in

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See the related Commentary beginning on page 656.
response to classic stimuli, such as LPS, is actually inhibited by LT treatment (20). This is not surprising, as this protease cleaves major components of three MAPK pathways (20–24). TNF receptor and IL-1 receptor knockout mice are not more resistant to spore challenge than their C57BL/6J parents (25), and LT-susceptible species such as rats (often used for testing of antitoxin therapeutics), as well as humans, harbor toxin-resistant macrophages (26).

Our study investigated the mechanism by which LT kills mice. BALB/c “sensitive” and C57BL/6J “resistant” mouse strains (harboring sensitive and resistant macrophages, respectively) were compared at the histopathological and molecular levels. We found a striking contrast between the two strains in early cytokine induction but ultimately similar pathology that did not involve macrophage susceptibility or TNF-α. Mortality in LT-treated mice was related to hypoxia-induced liver failure and pleural transudate.

Methods
Toxin. PA and LF were prepared and purified as previously described (27). Mutant LF687C has been described (28). Mutant PA678SA is a nonbinding PA mutant (M.J. Rosovitz, personal communication). Toxin was diluted in PBS and filter sterilized prior to use. Doses of LT refer to equal amounts of each component (i.e., 100 µg LT is 100 µg PA plus 100 µg LF). PA and LF were always injected together in the same PBS solution.

Animals. Eight- to ten-week-old male BALB/c and C57BL/6J mice were purchased from the Jackson Laboratory (Bar Harbor, Maine, USA). C57Bl/TnfSF6dld (gld) mice (in which mutated, nonfunctional FasL has been backcrossed for 15 generations onto a BALB/c background) were a generous gift from Thomas J. Sayers (National Cancer Institute, Frederick, Maryland, USA). Mice were injected with toxin dilutions, either 1.0 ml intraperitoneally (i.p.) or 10–100 µl intravenously (i.v.). Mice were euthanized by CO2 at various times after injection and terminally bled before necropsies were performed. Hematological parameters and differential leukocyte counts were analyzed by the NIH Clinical Center Department of Laboratory Medicine using the Cell-Dyne 3500 automated complete blood count (CBC) analyzer (Abbott Laboratories, Abbott Park, Illinois, USA), and serum was separated for cytokine analyses. Blood chemistry analyses for renal and hepatic function were performed using a Roche/Hitachi 917 analyzer (Roche Diagnostics, Indianapolis, Indiana, USA). Urine, collected at various time points after injection, was analyzed for urea nitrogen and creatinine using a Beckman Coulter Synchron LX20 instrument (Beckman Coulter Inc., Fullerton, California, USA).

Histopathology. A complete necropsy was performed on each mouse. Tissues were fixed by submersion in 10% neutral-buffered formalin, embedded in paraffin, stained with H&E, and evaluated by a board-certified veterinary pathologist. In the initial dose-finding experiments using 30, 50, 100, and 250 µg toxin (i.v. and i.p. routes), more than 40 tissues per animal were evaluated from several animals per dose. The i.v. and i.p. routes produced similar histopathological findings. Based on identification of the most consistently affected tissues, selected tissues were then evaluated in a timed sacrifice study using 100 µg LT (i.p.) in sets of three BALB/c and three C57BL/6J mice euthanized at 6, 12, 18, 24, 36, and 48 hours after injection. Control mice injected with PBS or PA alone (100 µg) were necropsied at 24 and 48 hours. Mandibular and mesenteric lymph nodes, liver, spleen, lung, heart, thymus, adrenals, and femur with marrow were evaluated. Select sections were stained by phosphotungstic acid hematoxylin (PTAH) or Masson’s trichrome. Severity of lesions was qualitatively graded on a four-point scale: minimal, mild, moderate, and marked. Immunohistochemistry was performed on select tissues: mandibular and mesenteric lymph nodes, thymus, liver, spleen, and lung. Antibodies to CD45R (Pharmingen, San Diego, California, USA), CD3 (DAKO Corp., Carpinteria, California, USA), and the macrophage surface antigen F4/80 (Caltag Laboratories Inc., Burlingame, California, USA) were used for formalin-fixed tissues. Vectastain Elite kits (Vector Laboratories Inc., Burlingame, California, USA) were used for stain visualization, and diaminobenzidine (DAB) was the chromagen. Antibodies against murine heat shock protein 70 (HSP70) (Santa Cruz Biotechnology Inc., Santa Cruz, California, USA), VEGF (Santa Cruz Biotechnology Inc.), and heme oxygenase-1 (HO-1) (StressGen Biotechnologies Corp., Victoria, British Columbia, Canada) were on 4% paraformaldehyde-fixed tissues. The EnVision kit (DAKO Corp.) with DAB was used. In separate experiments, spleens were extracted and immediately placed in RNAlater (Ambion Inc., Austin, Texas, USA) followed by RNA extraction using Trizol (Life Technologies Inc., Rockville, Maryland, USA), for use in the RNase protection assays (RPAs) described below. Alternatively, spleens were placed in 4% paraformaldehyde for in situ staining, described below. For Western blotting of tissue antigens, livers were harvested and homogenized in RIP buffer (1× PBS, 1% Nonidet P-40, 0.5% sodium deoxycholate, 0.1% SDS) using Tissue-Tearor (BioSpec Products Inc., Bartlesville, Oklahoma, USA). Homogenates were spun, and protein was quantified using the BCA kit (Pierce Chemical Co., Rockford, Illinois, USA) prior to SDS-PAGE and immunoblotting. Cells. RAW264.7 macrophage cells were cultured in DMEM with 10% FCS. Peritoneal macrophages were elicited by injection of 2 ml of 4% Brewer modified thioglycollate (Becton Dickinson Microbiology Systems, Cockeysville, Maryland, USA) 72 hours prior to harvesting by peritoneal lavage. Monocytes were extracted from mouse peritonea by lavage without prior thioglycollate injection. Mononuclear cells were attached to plastic and washed with medium prior to culture.

RPA. The Multiprobe RNase Protection Assay (RPA) (Pharmingen) was performed with some modification. Probes were synthesized with 33P-UTP (70–80 µCi per full reaction) using the Pharmingen In Vitro Transcrip-
and differences in i.p. and i.v. survival curves at both 50 and 100 µg dose injected i.p., presented to allow direct comparison of the single selected dose used in all further experiments.

While the number of circulating monocytes initially rose to about 200 bp. Following hybridization, slides were dipped in radiographic emulsion, exposed for 5 days, developed, and stained with H&E.

**Erythropoietin measurements.** The human erythropoietin (EPO) immunoassay kit (R&D Systems Inc.) can be used for mouse EPO measurements and was used to assess serum levels of EPO at various times after toxin injection.

**In situ hybridization.** In situ hybridization was performed by a previously published protocol originally developed for detection of viral RNA (29). Tissue sections were hybridized with 35S-labeled riboprobes in the sense and antisense configurations transcribed from a 1.0-kb KC gene insert. KC cDNA was cloned via RT-PCR into the Topo-TA cloning vector pCRII-TOPO (Invitrogen Corp., Carlsbad, California, USA). The plasmid was linearized before transcription with either T7 polymerase or SP6 polymerase. Probes were truncated by alkaline hydrolysis to about 200 bp. Following hybridization, slides were imaged in a Fuji 5000 PhosphorImager (Fuji Corporation, Tokyo, Japan). Subsequently, the slides were dipped in radiographic emulsion, exposed for 5 days, developed, and stained with H&E.

**Results**

**LT-induced lethality in inbred mice**

The effects of LT dose and route of administration were examined in BALB/cj and C57BL/6j mice, which have toxin-sensitive and -resistant macrophages, respectively (30). Mortality consistently occurred earlier and more often in the BALB/cj mice (Figure 1, a and b). Since both i.p. and i.v. routes produced similar survival curves at the 100-µg dose (Figure 1c), i.p. administration of this dose was used for subsequent experiments. At 100 µg i.p., BALB/cj mice displayed severe malaise by 48 hours, with 50% dying by 60 hours. By comparison, at 48 hours, C57BL/6j mice showed minimal to no malaise, and few of these mice died before 60 hours. However, 40% of C57BL/6j mice were dead by 96 hours (Figure 1d).
decreased in the C57BL/6J mice, there was rapid recovery to higher than initial levels after 10 hours (Figure 2a). Interestingly, the marked depletion of circulating mononuclear cells was not paralleled by depletion of fixed tissue macrophages (see below). There was a rapid five- to tenfold increase in circulating neutrophils, with numbers remaining high over 48 hours in both strains (Figure 2b). A drop in platelet levels (thrombocytopenia) occurred in both strains in the first 12 hours and was more marked in the BALB/cJ mice (Figure 2c). However, there were large variations in platelet numbers in all groups harvested at 24 hours or longer after LT administration, and many terminally ill animals did not display thrombocytopenia. Thus, thrombocytopenia was not consistently associated with terminal illness. There were no changes in circulating B and T lymphocyte numbers (data not shown).

Histopathological and physiological findings in LT-treated mice

Summary of macroscopic findings. These findings are cumulative from multiple experiments with a total pool of six to twelve animals per time point. No significant gross lesions were observed in control mice. Significant observations in the BALB/cJ mice included the following: multiple tan or dark red liver foci in 90% of the mice at 18 hours and later time points; mottled “nutmeg” liver in 14 of 30 at 60 hours; discolored dark red/black feces in the small and large intestines in 50% at 18 hours and later time points; dark splenic foci, only at 18 hours; darkened paws and black toes in 19 of 60 at time points past 12 hours; pleural cavity fluid (1–3 ml) in 60% at 48 hours and in 90% by 60 hours; and ascites (i.e., peritoneal cavity fluid; 3–5 ml) in 50% at 60 hours. C57BL/6J mice developed similar lesions but at later times: tan to red liver foci in 5% of animals at 48 hours and 50% by 72 hours; mottled “nutmeg” liver in 12 of 30 at 96 hours; discolored feces in 20% at 96 hours; pleural cavity fluid in 90% at 96 hours; and ascites in 30% at 96 hours. Darkened extremities and splenic foci were not seen in the C57BL/6J mice.

Microscopic findings and chemical analyses. No significant lesions were noted in the control mice. In mice receiving LT, lesions occurred primarily in the spleen, bone marrow, and liver, with more minor changes in a few other tissues, as described below.

Spleen. At 6 and 12 hours after injection, BALB/cJ mice treated with LT showed mild to moderate individual cell (apoptotic-like) death of myelogenic cells in the splenic red pulp (Figure 3, b and c) accompanied by a moderate decrease in normal extramedullary hematopoiesis (EMH), including megakaryocytes. Minimal to mild individual (apoptotic-like) death of lymphoid cells was present in the splenic white pulp. At 18 hours, mild to moderate hemorrhage developed in both the red and the white pulp (Figure 3, d and e). A turnaround occurred by 24–36 hours and continued through 48 hours (Figure 3f), with a decrease in splenic necrosis, resolution of the hemorrhage, and recovery of EMH with a left shift (increase in percentage of immature versus mature forms, primarily in the granulocytic series). The C57BL/6J mice, in contrast, showed no...
changes in the spleen, with the exception of a minimal increase in EMH only at 18 hours (data not shown).

**Bone marrow.** At 6 and 12 hours after injection, BALB/cJ mice showed mild to moderate individual cell death throughout the femoral bone marrow (Figure 4b). Severity of the apoptotic-like bodies peaked at 18 hours (Figure 4c). By 24 hours, mild, diffuse, infarction-like necrosis, compatible with hypoxia, was evident in the metaphyseal marrow of the femur. Neither the diaphyseal nor the epiphyseal marrow showed this overt necrosis. By 48 hours, necrosis in the metaphyseal bone marrow was marked (Figure 4, e and f). Individual cell death in the diaphyseal marrow was also noted to a mild degree. The C57BL/6J mice showed similar pathologies in the bone marrow at later time points and with lower incidence and severity. Mild metaphyseal marrow necrosis first occurred at 36 hours and became moderate at 48 hours in the C57BL/6J mice (data not shown).

**Liver.** BALB/cJ mice developed liver lesions ranging from small hemorrhagic infarcts to large areas of centrilobular coagulative necrosis by 18–24 hours after LT injection (Figure 5b). Rarely, mild to moderately dilated vascular spaces with poorly formed acellular thrombi were associated with the necrosis (Figure 5c). No significant amount of fibrin was evident in these poorly organized thrombi (PTAH stain) (data not shown). Liver necrosis increased in incidence and severity at 36 hours and continued with no recovery through 48 hours (Figure 5d). The presence of hemorrhagic infarcts and centrilobular coagulative necrosis without significant fibrin clots was compatible with hypoxia due to blood stasis and/or decrease in venous drainage rather than due to arterial thrombosis. Similar liver necrosis was noted at a later time point (48 hours) with lower severity (minimal to mild) in the C57BL/6J mice.

Severity progressed to marked by 72 hours and 96 hours in the C57BL/6J mice (data not shown). Analysis of blood chemistry revealed significant increases in the liver-dysfunction indicators serum alanine aminotransferase (ALT) and aspartate aminotransferase (AST) in a majority of the LT-treated mice at time points after 48 hours (Figure 6, a–d). Liver dysfunction was also indicated by decreased serum albumin levels noted in both mouse strains in response to toxin over time (Figure 6, e and f).

**Thymus and heart.** The BALB/cJ mice had minimal lymphoid cell death in the thymic cortex and medulla at 6 hours, without significant change through 24 hours. At 36 hours, T lymphocyte death was moderate (probably secondary to the stress of major organ necrosis/failure), progressing to marked levels at 48 hours, with a resultant decrease in cortical depth (data not shown). Minimal coagulative necrosis was noted in the heart, with occasional mild cardiac dilatation at 48 hours in a few animals (data not shown). Blood chemistry analysis, however, indicated that the serum creatinine kinase level (an indicator of myocardial dysfunction) was elevated up to fivefold in only 10% of toxin-treated mice of both strains at later time points (data not shown).

**Other observations.** Extensive pleural and peritoneal fluid accumulation with progression of malaise state was noted in both mouse strains. Despite the relatively high incidence of pleural fluid accumulation noted grossly, light microscopy did not reveal any lesion in the lung, suggesting that permeability alterations did not result from overt tissue damage. Neither H&E nor PTAH staining of lung revealed any evidence of capillary thrombosis or fibrin clots as seen with the disseminat-
significant change in CRE was seen in either mouse strain at any time after LT treatment, with only 2 of 55 mice falling outside the normal 0.1–1.1 mg/dl serum CRE range for mice (Supplemental Figure 1; www.jci.org/cgi/content/full/112/5/670/DC1). Overall results indicated that there were no statistically significant changes in BUN levels in LT-treated mice, despite a small number of outliers (3 of 27 BALB/cJ and 4 of 28 C57BL/6J mice) (Supplemental Figure 1). These mice may not represent biologically relevant events, since BUN changes independent of CRE are not an indication of renal dysfunction and often reflect altered metabolism or urination. No significant changes in the levels of urea nitrogen and CRE were seen in urine from LT-treated animals collected every 12 hours after injection (data not shown).

There was no evidence of neutrophil margination in the blood vessels of any examined tissue. Even areas of extensive tissue necrosis, such as the liver, had minimal to no associated inflammatory cells. Immunohistochemical staining for macrophages showed only

ed intravascular coagulation (DIC) that is commonly associated with classical macrophage/TNF-α-driven endotoxic shock. Because hypoalbuminemia caused by liver damage can directly lead to leakage of fluid from vessels due to low plasma colloid osmotic pressure, we hypothesized that this effect might contribute to the vascular leakage observed in the LT-treated mice. Analysis of peritoneal and pleural fluids from premortem mice revealed cell-free, clear, straw-colored fluids with lower albumin concentrations relative to serum (Figure 6, e and f). This albumin ratio is used to differentiate between exudates resulting from inflammation-induced increased capillary permeability and transudates resulting from altered intravascular hydrostatic/oncotic pressure. The fluids in LT-treated mice appear to be the latter, supporting a contribution of low colloid osmotic pressure to the fluid accumulation seen in LT-treated animals.

H&E and PTAH staining of kidneys, the other organ often affected in cytokine-mediated shock, did not reveal any lesions, nor evidence of capillary thrombosis. Evidence that renal failure did not occur in LT-treated mice was further provided by analysis of blood urea nitrogen (BUN) and serum creatinine (CRE). Serum CRE is the more specific determinant of renal function, because, unlike BUN levels, serum CRE levels are not affected by protein metabolism or the hydration status. No significant change in CRE was seen in either mouse strain at any time after LT treatment, with only 2 of 55 mice falling outside the normal 0.1–1.1 mg/dl serum CRE range for mice (Supplemental Figure 1; www.jci.org/cgi/content/full/112/5/670/DC1). Overall results indicated that there were no statistically significant changes in BUN levels in LT-treated mice, despite a small number of outliers (3 of 27 BALB/cJ and 4 of 28 C57BL/6J mice) (Supplemental Figure 1). These mice may not represent biologically relevant events, since BUN changes independent of CRE are not an indication of renal dysfunction and often reflect altered metabolism or urination. No significant changes in the levels of urea nitrogen and CRE were seen in urine from LT-treated animals collected every 12 hours after injection (data not shown).

There was no evidence of neutrophil margination in the blood vessels of any examined tissue. Even areas of extensive tissue necrosis, such as the liver, had minimal to no associated inflammatory cells. Immunohistochemical staining for macrophages showed only
a minimal decrease in fixed-tissue macrophages in the lymph nodes, spleen, and liver at the various time points (Supplemental Figure 2; www.jci.org/cgi/content/full/112/5/670/DC1), indicating that the effect of LT on these cells was different from its effect on circulating monocytes. Immunohistochemistry also showed minimal to no decrease in T cell populations in spleen, liver, and lung (data not shown). A minimal to mild increase in B cells was noted, particularly at later time points in the liver and lung (data not shown). Despite the presence of red intestinal contents in some mice, there was no evidence of intestinal hemorrhage or necrosis by light microscopy. Hemorrhage (often associated with DIC) was also not noted in any other organ, with the exception of the BALB/cJ spleens at 18 hours. Limbs macroscopically noted to be discolored exhibited congestion and edema, without evidence of thrombi or inflammation.

In summary, crisis in the mice appeared to relate to liver necrosis resulting from hypoxia. Oxygenation insufficiency and systemic hypoxia may be the result of vascular permeability alterations, reflected by the accumulation of pleural and peritoneal cavity fluid and gross edema in extremities, or, alternatively, may be the initiating factor for the observed vascular leakage. Metaphyseal bone marrow necrosis observed in LT-treated mice also supports tissue injury due to oxygenation insufficiency.

LT induction of hypoxia

To further investigate LT-induced hypoxic injury, EPO, HO-1, and HSP70 were used as markers of hypoxia (31–34). Induction of these genes by a similar stimulus may differ based on mouse-strain differences. Therefore, response levels do not provide a quantitative measure of illness, extent of vascular collapse, or progression of hypoxic state between different mouse strains. Their induction, however, is a clear indicator of a hypoxic response. Analysis of sera from LT-treated mice showed remarkable increases in EPO in both strains (Figure 7a), which may reflect systemic hypoxia. These data also support the previous observation that kidneys are not affected in LT-treated mice, since the kidney is a major source of EPO, and renal dysfunction manifests as a decrease in EPO production. Immunostaining of liver showed a slight increase in HO-1 in hepatocytes and a marked increase in Kupffer cells at 48 hours in both strains (Supplemental Figure 3). Western blot analysis for HSP70 revealed an impressive induction in the liver (Figure 7b). These results demonstrate a hypoxic state induced in mice by LT treatment.

LT induction of cytokines and chemokines in mice

To investigate the potential involvement of cytokines in LT-mediated pathogenesis (18), the production of numerous factors was screened for by RPA analysis: IL-4, IL-5, IL-3, IL-13, IL-9, IL-2, IL-12, IL-10, IL-1α, IL-1β, IL-1RA, IL-18, IL-6, MIF, Ltn, etoxin, RANTES, MIP-1B, MIP-2, MCP-1/JE, IP10, C10L, KC, LIF, IL-7, M-CSF, SCF, IFN-γ, FasL, caspase-8, Fas, FADD, FAP, FAI, TRAIL, TNFRp55, TRADD, RIP, TNF-β, LTB, TNF-α, and TGF-β1. Three splens per

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**Figure 7**

LT-induced hypoxia. (a) Mice were injected i.p. with 1.0 ml of PBS or 100 µg LT. Serum EPO measurements at each time point represent averages of values assessed for two samples, with each sample representing sera pooled from three mice. The exception is the 72-hour time point for BALB/cj, which represents measurement of one sample composed of pooled sera from three mice. (b) Liver extracts from BALB/cj or C57BL/6j mice treated i.p. with 100 µg LT for 6, 24, 60, and 72 hours were analyzed for HSP70 levels by Western blot.

**Figure 8**

RPA analysis of LT-treated mice. (a) BALB/cj mice were treated i.p. with 100 µg LT, and spleens were obtained after 10 or 32 hours. PBS-injected and noninjected (NI) controls were included for comparison. (b) A similar experiment with additional time points. A PBS-injected control was included for comparison. (c) Comparison of BALB/cj and C57BL/6j spleen KC RNA levels at various times after i.p. injection of 100 µg LT. GADPH was used as a control for equal RNA loading in all lanes.
time point were harvested from 2.5 hours to 72 hours (BALB/c) or 2.5 hours to 96 hours (C57BL/6J) after LT injection. Cytokine and chemokine expression was not altered between 24 hours and 96 hours; however, a number of factors were induced between 2.5 and 24 hours, primarily in the BALB/cJ mouse. Representative RPA gels are shown in Figure 8, and data from these and other gels are summarized in Table 1. Factors that showed altered expression in the BALB/cJ mouse, as early as 2.5 hours, included IL-10, GM-CSF, IL-6, eotaxin, MIP-2, MCP-1/JE, KC, LIF, and M-CSF (Figure 8, a and b). The induction of the KC gene was notably rapid and dramatic in the BALB/cJ spleen samples (Figure 8c). The C57BL/6J mouse did not show alterations in the majority of these factors, with the exception of a slight early induction of KC and M-CSF. IL-15 was induced only in the C57BL/6J mouse (Table 1). Although previous studies have reported production of IL-1β and TNF-α by macrophages and implicated these cytokines in an endotoxic shock–like death mechanism (18), our analysis showed no induction of IL-1β or TNF-α at the RNA level. In the earlier studies, done in cultured macrophages, LT preparations were produced in Escherichia coli, and trace amounts of endotoxin may have been responsible for the cytokine response. Our findings detected no evidence of classic Th1/Th2 cytokine responses or TNF-α–based inflammatory cascades in either strain. Analyses performed on spleens from mice injected by the i.v. route or with lower toxin doses gave similar results (data not shown). Factors listed above that do not appear in Table 1 showed no changes at the RNA level.

Protein profiles analyzed from sera by ELISA (Figure 9) closely paralleled RPA results from spleen, with a few exceptions. Antinflammatory IL-10 and proinflammatory LIF, which showed moderate RNA induction (Figure 8, a and b), were not detected at the protein level (data not shown). Also, IL-1β, which was not induced at the RNA level in spleen, was detected in serum of toxin-treated BALB/cJ mice. IL-1β may have been induced in other organs, or simply released from intracellular stores. The C57BL/6J mice showed only a slight KC response, which was also noted at the RNA level (Figure 8c). By 24 hours, all factors except MCP-1/JE and G-CSF had declined to baseline levels. These latter two factors returned to base line by 48 hours. The induction of TNF-α was not observed in either strain at the protein level (Figure 9).

To test specificity of cytokine induction, additional mice were injected i.p. with 100 µg of PA, LF, mutant PA (D683A substitution) that does not bind to receptor (M.J. Rosovitz, personal communication), mutant LF (E687C substitution) that does not have proteolytic activity (28), a combination of mutant PA and mutant LF, boiled LF, or BSA. Both PA and LF induced KC, MCP-1/JE, and G-CSF transiently at 2.5 hours, but at 25–50% of the LT-induced levels, and induction was at or below that in PBS-treated controls by 6 hours (data not shown). Since BSA, mutant toxins, and boiled LF also induced these three factors, induction was interpreted to be a nonspecific transitory response to protein (not PBS). None of the other cytokines was induced by PA or LF (data not shown). The significantly higher induction and extended course of response beyond 6 hours in LT-treated mice indicate that induction is specific to active toxin in the BALB/cJ mice.

**LT effects on VEGF expression**

VEGF has been associated with increased vascular permeability in the lungs (35, 36) and other organs (37) and is induced by hypoxia (38). Pleural effusions associated with mortality in both mouse strains led to careful examination of this cytokine. VEGF was not induced in liver, spleen, or lung tissues of LT-treated mice analyzed by RPA and Western blot. Immunostaining of lungs also revealed no local increase in VEGF expression (data not shown).

**Role of FasL in LT-mediated mortality and hepatic injury in mice**

Hepatocytes are sensitive to programmed cell death, and Fas/FasL apoptosis pathways play a major role in liver pathology in numerous diseases (39). Polymorphisms between C57BL/6 and BALB/c mice affect specific activity of FasL, with higher cytotoxic function associated with the BALB/c strain (40). Although LT-induced FasL (Table 1) and soluble FasL were measurable in the sera of BALB/cJ mice (Figure 10a), LT-treated FasL-deficient (gld) mice exhibited a mortality
curve (Figure 10b) and pleural cavity fluid accumulation (data not shown) similar to those of their BALB/cJ counterparts, indicating that LT cytotoxicity does not require FasL function. Lack of LT-mediated FasL induction in C57BL/6J mice supports this result (data not shown).

LT induction of cytokines from isolated macrophages

Previous studies have reported cytokine production from macrophages in response to sub-lytic doses of LT (18, 41, 42). BALB/cJ macrophages are extremely sensitive, with 100% of cells dying 90–120 minutes after treatment with 200 ng/ml of LT, while C57BL/6J macrophages are resistant, with 100% surviving at 6 hours after treatment with 2,000 ng/ml toxin (data not shown). BALB/cJ peritoneal macrophages and RAW264.7 cells (a sensitive macrophage cell line) treated with 10 pg/ml (sub-lytic) to 1,000 ng/ml (lytic) of LT for 90 minutes, 120 minutes, or 6 hours, as well as culture supernatants, were tested for production of the cytokines shown in Figure 9. RAW264.7 cells (with or without toxin treatment) constitutively produced a high level of MCP-1/JE (4,500 pg/ml), which was not altered by LT treatment. None of the other factors was induced by LT treatment of in vitro macrophages from either mouse strain. To identify a cellular source in vivo, we performed in situ analysis of KC expression and confirmed the impressive induction of this chemokine in spleen and liver of LT-treated BALB/cJ mice at 2.5 hours (Figure 11a) and 6 hours (Figure 11b). No expression remained at 24 hours (Figure 11c), and no expression was seen in PBS-injected controls (data not shown). By comparison, minimal KC was evident at 6 hours in C57BL/6J mice (Figure 11d). The diffuse high expression seen throughout the liver of the BALB/cJ mice at 2.5 hours and 6 hours was associated with hepatocytes and not Kupffer or endothelial cells (Figure 11e). KC expression decreased when hepatic necrosis began (data not shown). Autoradiographic analysis of the spleen sections (Figure 11, f–h) revealed KC expression in the marginal zone, follicular dendritic cells, and red pulp. LT induction of KC in hepatocytes at such early time points, prior to appearance of hypoxia or liver damage, presents, to our knowledge, the first report of the toxin directly affecting this cell type and introduces a potentially new direct target for LT-mediated effects.

**Figure 9**

Time course of serum cytokine protein levels after LT treatment. Mice were injected i.p. with 1.0 ml of PBS or LT (TOX; 100 µg PA + 100 µg LF). ELISA determinations were done on sera pooled from three to five mice for each time point and treatment.
Discussion
In this study, events leading to LT-induced death of BALB/cJ and C57BL/6J mouse strains were analyzed in detail. Despite harboring toxin-sensitive and resistant macrophages, both strains are susceptible to anthrax infection (17, 25) and, as reported here, to LT. BALB/cJ mice succumb more rapidly, but both strains experience a similar course of disease, with hypoxia-mediated necrosis of liver and metaphyseal bone marrow as hallmarks of crisis, accompanied by pleural effusions. Contrary to previous hypotheses (18), hallmarks of cytokine-induced shock such as inflammation, fibrin clots and associated thrombosis, DIC, neutrophil margination and associated endothelial cell damage, renal dysfunction, and TNF-α production are absent in LT-treated mice. Despite an early transitory production of other cytokines in BALB/cJ, no inflammatory cascade develops. C57BL/6J, which do not mount this response, display similar although delayed pathology and subsequent death. The data indicate that LT induces hypoxia-mediated tissue damage and cytokine-independent death with shocklike manifestations. Macrophage lysis is not essential for LT-mediated death.

The cause of the more rapid onset of pathology in the BALB/cJ mice is unclear. The rapid lysis of their macrophages with accompanying cytokine response may increase the rate of LT-mediated events but is clearly not required for the hypoxia, liver necrosis, pleural effusions, and mortality seen in both strains. The cytokine burst is not necessarily linked to macrophage lysis, although the rapid release of IL-1β, as early as 150 minutes, probably reflects release of intracellular stores concomitant with lysis, since induction of IL-1β mRNA is not observed. Transitory production of other cytokines could be a response of various cells to IL-1β (43–47). The remarkable production of eotaxin and the KC induction in hepatocytes confirm involvement of other cell types and at the same

Figure 10
LT induction of FasL and cytotoxicity in FasL-deficient mice. (a) BALB/cJ mice were treated i.p. with 100 µg LT, and ELISA determinations were done on sera pooled from three mice for each time point. (b) BALB/cJ (n = 6) or FasL-deficient (n = 6) mice were injected i.p. with 100 µg LT, and time to death was recorded for each strain.

Figure 11
Induction of KC in spleen and liver analyzed by in situ hybridization. BALB/cJ or C57BL/6J mice were injected i.p. with 100 µg LT. Spleens and livers were harvested at 2.5, 6, 24, or 36 hours after injection for in situ analysis of KC production. (a–c) Whole-organ-section phosphorimages of sense (s; control) and antisense (as) 35S KC probe hybridization to BALB/cJ spleen and liver at 2.5 hours (a), 6 hours (b), and 24 hours (c). (d) Sections of C57BL/6J mouse organs at 6 hours after LT. Arrows in a–d indicate the single spleen section in each panel; all other sections are liver. Red signifies the highest intensity of signal, while bluish-green is negative background. (e–h) Silver-grain densities in various regions of each organ. (e) High clustering of grains at liver hepatocytes, with little association with endothelial cells (EC). (f) Higher expression in the spleen marginal zone (MZ) in contrast to white pulp (W). (g) A dark-field image similarly showing the higher silver-grain densities associated with marginal zone (MZ), red pulp (R), and follicular dendritic cells (FDC). (h) High density of silver grains at the FDCs of the white pulp.
time support a requirement for macrophage lysis, since hepatocytes are expected to be equally sensitive in both mice. C57BL/6J macrophages are not defective in normal cytokine responses to other insults (48, 49).

Immunological differences between BALB/c and C57BL/6 in sensitivity and resistance to a variety of bacterial, parasitic, and fungal infections have often been linked to Th1/Th2 responses (50–52). However, classic Th1/Th2 cytokines, and those involved in their regulation (IL-12, IL-4, IFN-γ, etc.), are not induced by LT, nor are inflammation or traditional responses associated with IL-1β and IL-6 observed. It is therefore difficult to link the exacerbation of LT toxicity in BALB/cj mice to these or other induced cytokines (MCP-1/JE, eotaxin, GM-CSF, etc.). Strain differences in FasL (40) and NO production by macrophages (53, 54) are also not involved in LT susceptibility (experiments with iNOS knockout mice; data not shown). Ultimately the difference between these strains may lie not in the brief cytokine response seen in BALB/cj, but in the induction of a protective response in C57BL/6J, such as the LT-mediated IL-15 induction.

The difference in LT susceptibility between the two strains may not be linked to immune responses. Potential strain differences might include the number of toxin receptors. Although macrophages from both strains internalized LT and showed cleavage of MAPKK proteins with similar kinetics (data not shown), it is not known whether an equivalent number of toxin receptors exist on all tissues. Additional genetic differences could result in differential resistance at a variety of levels, ranging from neuroendocrine control of vascular barrier function to general stress response by undefined molecular mechanisms. Documented differences in the response of these two strains to neural and hormonal elements (55, 56) could have consequences for pathological cascades. These differences need not correlate with downstream effects on cytokine function (57). The higher resistance of C57BL/6 mice to hypoxic insult may provide an even simpler explanation for their delayed death (58, 59).

Regardless of the mechanism of cytokine response in LT-treated BALB/cJ mice, and its possible role in pathology in this strain, neither macrophage lysis nor this response is required for LT-induced lethality in mice. Our findings confirm that LT does not induce TNF-α in LT-sensitive mouse macrophages (19). The lack of a sustained cytokine response is consistent with LT-mediated shutdown of MAPK signaling (20, 21). Despite the extensive emphasis given to the role of macrophage lysis in LT effects in anthrax (18), the lack of a requirement for macrophage lysis is not surprising, given that lysis occurs only in a limited subset of inbred mice. Human, rat, and many LT-sensitive mouse strain macrophages are resistant to LT (data not shown).

Death induced by LT in mice may relate to liver necrosis accompanied by pleural effusions. Hypoxia is clearly involved in the tissue necrosis. The histopathological observations in LT-treated mice, including hepatic centrilobular coagulative necrosis, metaphyseal bone marrow necrosis, and infarction-like necrosis of the incisor dental pulp (data not shown), are indicative of hypoxic tissue death. Hypoxia-marker analyses at the organ level and high levels of EPO in blood confirm LT-induced hypoxia. Hypoxia and tissue damage are accompanied by pleural effusion, ascites, and congestion and edema of the extremities, all generic indicators of circulatory collapse. Whether hypoxia-linked tissue necrosis is the primary event leading to a secondary vascular leakage and shocklike symptoms or is itself a result of circulatory insufficiency due to LT effects on vascular function is unknown.

Vascular dysfunction and thrombosis have been described in association with anthrax infection in humans (9, 60) and animals (60–64). This type of classic vasculitis induced by endothelial adherence of inflammatory cells and fibrinoid necrosis of vessels did not occur in LT-treated mice, despite similar accumulation of pleural fluid in human anthrax subjects, animal models (2, 9, 11, 14, 65, 66), and LT-treated mice. LT induction of shock by interference with endothelial cell function has been proposed (2, 7, 63, 64), but no evidence of direct LT effects on endothelialium has been presented. If vascular collapse occurs in LT-treated mice, it is quite different from that associated with endotoxic shock. Thrombocytopenia is present in 50% of LT-treated mice without evidence of thrombosis. Although often associated with consumptive coagulative dysfunction (i.e., DIC), thrombocytopenia may also be caused by direct platelet destruction, megakaryocyte loss or dysfunction, or hypoxia. Drugs can induce rapid thrombocytopenia without initial coagulative pathologies (67, 68). Additionally, there are examples of thrombosis-independent shock processes, such as the Hantavirus pulmonary syndrome (HPS), which manifests thrombocytopenia without coagulopathy. The pathology of HPS is intriguingly similar to that of LT: pulmonary effusions without histological changes in the lungs, neutrophilia, hypoalbuminemia, increased AST or ALT, no changes in renal or cardiac function, and no coagulation, but striking vascular leakage and multiorgan failure (69, 70).

An alternative hypothesis of LT-mediated toxicity is that LT induces hypoxia directly and the observed vascular leakage is secondary to tissue necrosis, rather than the cause of it. LT may interfere with metabolic pathways that prevent proper delivery of oxygen, or its utilization in tissues. As a consequence, extensive liver necrosis and subsequent hypoalbuminemia, with resulting decreased oncotic pressure, may drive the transudative pleural effusion seen in LT-treated mice. The later appearance of pleural fluid in LT-treated mice supports this hypothesis.

LT-treated mice show marked neutrophilia with no recruitment of these cells into any tissues. High neutrophil numbers have been previously reported with respiratory anthrax (3, 5, 11, 14, 71, 72) and are often associated with bacterial infections, but LT injection...
alone appears to induce massive release of the potent neutrophil-recruitment and -activation mediators MIP-2 and KC (which are similar to the IL-8 and GRO chemokines in humans). Exposure of neutrophils to LT has been shown to increase their chemotactic response to other substances (73). LT may also prevent proper passage of activated recruited neutrophils into tissues. A previous report indicates direct LT interference with P-selectin expression on BALB/c platelets in vitro (74). Preliminary studies indicate that LT induces the shedding of selectin molecules into the circulation (data not shown), and this could negatively impact neutrophil adhesion and traversal. Intravascular aggregation of activated neutrophils with subsequent release of reactive oxygen species and potent enzymes may play a role in tissue damage (75).

The detailed analysis of LT-mediated toxicity in mice presented here advances our understanding of anthrax pathogenesis. The similarities in the pathology induced by LT and by anthrax infection point to the importance of understanding how the toxin acts during infection. Further deciphering of the molecular events by which the toxin contributes to pathogenesis is a crucial next step in the study of anthrax. The mechanism linking LT proteolytic function to the in vivo pathogenesis that involves hypoxia and vascular dysfunction remains to be elucidated.

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